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Peltoniemi, Mikko

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# **Networked web-cameras monitor congruent seasonal development of birches with phenological field observations**

Mikko Peltoniemi<sup>1,\*</sup>, Mika Aurela<sup>2</sup>, Kristin Böttcher<sup>3</sup>, Pasi Kolari<sup>4</sup>, John Loehr<sup>5</sup>, Tatu Hokkanen<sup>1</sup>, Jouni Karhu<sup>1</sup>, Maiju Linkosalmi<sup>2</sup>, Cemal Melih Tanis<sup>2</sup>, Sari Metsämäki<sup>3</sup>, Jukka-Pekka Tuovinen<sup>2</sup>, Timo Vesala<sup>4</sup>, Ali Nadir Arslan<sup>2</sup>

1 Natural Resources Institute Finland (Luke), Latokartanonkaari 9, FIN-00790, Helsinki, Finland

2 Finnish Meteorological Institute, Erik Palménin aukio 1, FI-00560,

3 Finnish Environment Institute (SYKE), Mechelininkatu 34a, FIN-00251 Helsinki, Finland

4 Department of Physics, PO Box 68, 00014 University of Helsinki

5 Lammi Biological Station, University of Helsinki, Pääjärventie 320, 16900 Lammi, Finland

\*Corresponding author: Mikko Peltoniemi, mikko.peltoniemi@luke.fi, tel. + 358 40 801 5329,  
Latokartanonkaari 9, FIN-00790, Helsinki, Finland

## Abstract

Ecosystems' potential to provide services, e.g. to sequester carbon, is largely driven by the phenological cycle of vegetation. Timing of phenological events is required for understanding and predicting the influence of climate change on ecosystems and to support analyses of ecosystem functioning. Analyses of conventional camera time series mounted near vegetation has been suggested as a means of monitoring phenological events and supporting wider monitoring of phenological cycle of biomes that is frequently done with satellite earth observation (EO). Especially in the boreal biome, sparsely scattered deciduous trees amongst conifer-dominant forests pose a problem for EO techniques as species phenological signal mix, and render EO data difficult to interpret. Therefore, deriving phenological information from on the ground measurements would provide valuable reference data for earth observed phenology products in a larger scale. Keeping this in mind, we established a network of digital cameras for automated monitoring of phenological activity of vegetation in the boreal ecosystems of Finland. Cameras were mounted at 14 sites, each site having 1-3 cameras. In this study, we used data from 12 sites to investigate how well networked cameras can detect the phenological development of birches (*Betula spp.*) along a latitudinal gradient. Birches typically appear in small quantities within the dominant species. We tested whether the small, scattered birch image elements allow a reliable extraction of colour indices and the temporal changes therein. We compared automatically derived phenological dates from these birch image elements both to visually determined dates from the same image time series and to independent observations recorded in the phenological monitoring network covering the same region. Automatically extracted season start dates, which were based on the change of green colour fraction in spring, corresponded well with the visually interpreted start of the season, and also to the budburst dates observed in the field. Red colour fraction turned out to be superior to the green colour-based indices in predicting leaf yellowing and fall. The latitudinal gradients derived using automated phenological date extraction corresponded well with the gradients estimated from the

43 phenological field observations. We conclude that small and scattered birch image elements allow  
44 reliable extraction of key phenological dates for the season start and end of deciduous species studied  
45 here, providing thus important species-specific data for model validation and for explaining the  
46 temporal variation in EO phenology products.

47

48 Keywords: birch, budburst, camera, monitoring, phenology, time lapse

49

## 1 Introduction

Timing of spring onset has advanced significantly during the last century (Menzel et al., 1999, Menzel et al., 2006, Delbart et al., 2008, Jeong et al., 2011, Zhao et al., 2015). Seasonal variation of vegetation activity directly affects photosynthesis, growth of trees and plant reproductive investment, so it is an important driver of the global carbon balance and thus is strongly linked to climate change (Hogg et al., 2000, Richardson et al., 2013). A recent study that compared phenological data to predictions of 36 tree phenology models showed that both inter-annual and spatial variations of phenology is poorly predicted by the models (Basler et al., 2016). This is critical as the year-to-year variation in the timing of budburst of birches (*Betula* spp.) in the boreal zone varies in a wide range of 40 days (Häkkinen, 1999). Poor reproduction of the phenological cycle in biosphere models has also been shown to cause a consistent overestimation of carbon balance in comparison to measured data (Richardson et al., 2012, 2013). The predictive power of models can be expected to further degrade under climate change, due to decoupling of light and temperature cycles. Decoupling of these cycles will be pronounced in northern latitude forests, which are expected to face increases of mean temperatures by 2-7 °C (Ruosteenoja et al., 2016). Therefore, continuous, long-term monitoring of vegetation activity is needed.

Phenological monitoring has a long tradition, and phenological observation networks exist in many countries across the world (Siljamo et al., 2008). At the same time, many spectro- and radiometric instruments suitable for phenological monitoring are operating from space, complementing the dating of phenological events over wider regions (Zhang et al. 2006, Böttcher et al., 2014, Gonsamo et al., 2016). In recent years, also near-surface remote sensing with time lapse imaging (Richardson et al., 2007) has provided a cost-effective methodology to monitor and ground-truth phenological phenomena (Hufkens et al., 2012, Klosterman et al., 2014). Time lapse imaging solves some of the problems associated with traditional field observations, as more quantitative methods can be used to define the start of the growing season, for example, while still maintaining the link to the visual appearance of

plants. Time-lapse image based phenological development could also provide a closer analogy to remote sensing than field observations of phenology, which are not fully comparable with remote sensing observations as they detect different traits (Badeck et al., 2004). Methodologically, automated curve fitting and transition date extraction methods used for camera image time series have similarities with EO data processing (Elmore et al., 2014; Klosterman et al., 2014).

Cameras have most often been used to analyse the phenological development of deciduous species (Richardson et al., 2007), although also other types of ecosystems, such as grasslands (e.g. Migliavacca et al., 2011), peatlands (Westergaard-Nielsen et al., 2013, Peichl et al., 2015, Linkosalmi et al., 2016) and coniferous forests (Nagai et al., 2012, Linkosalmi et al., 2016), have been monitored. Analyses are robust to the scene illumination angle, cloud cover and camera type, if suitable analysis methods are used (Sonnentag et al., 2012; Linkosalmi et al., 2016; Peltoniemi et al., 2017). Colour changes in plant tissue are unlikely to occur without a biochemical or biophysical mechanism, and digital photography has provided insight into these mechanisms (Keenan et al., 2014; Yang et al., 2014). For deciduous species, budburst and leaf senescence events and also their relationship with CO<sub>2</sub> exchange have been in a focus in a number of studies, and these phenomena have been analysed with various colour indices (Richardson et al., 2007, Ahrends et al., 2009, Sonnentag et al., 2012, Mizunuma et al., 2013, Wingate et al., 2015).

There are still open questions regarding how the camera-derived phenological data should be used in an optimal way. It would be interesting to know how the image-extracted dates compare with those based on the field definitions used in phenological observation networks, and which transition dates can be extracted with sufficient accuracy. This would provide more solid basis for using cameras to supplement existing field observation networks. Secondly, a single image may provide a wealth of information on several species, some only appearing in the margins of the image or amidst the dominant vegetation in

smaller proportions and the understory, but the use of such information has been rare. Still, the non-dominant elements potentially provide important information for interpreting earth observations, which aggregate information from an area that may not be fully represented by the species dominating the camera view (Hufkens et al., 2012). In the boreal zone, deciduous trees often occur in relatively small and fragmented areal proportions in the satellite footprint. While the areal proportion may be small, their phenology causes distinctive changes in the reflective properties of canopies (Böttcher et al., 2014; Jönsson et al., 2010), which complicates phenological analyses of conifers, and may even render results unreliable.. Species-specific phenological information drawn from image time series combined with high-resolution earth observation data on species distributions could markedly improve the quality of satellite-based phenology estimation (Liang et al., 2011; Liu et al.2015). If part of the monitoring would be based on scattered and smaller species-specific image elements, and not only camera views dominated by species, the cost of representative monitoring of wide area phenology would naturally be reduced.

We established a network of cameras at 14 boreal sites in Finland, each including 1-3 cameras in different positions. Most of the sites in the network are dominated by Scots pine (*Pinus sylvestris*) and Norway spruce (*Picea abies*), and some are peatlands. Twelve of the sites have a varying mixture of *Betula spp.*, allowing a cross-site study of their phenology, and making it possible to study how these sometimes small and marginal image elements of widely distributed species could benefit phenological monitoring using web-cameras.

The objectives of this study were to test the use of the recently established camera network for birch phenology analysis and supplementing existing phenological field observations. We were interested in how selected color indices compare to the conventional phenological observations, and whether the scattered and often small birch elements within the images provide a useful source of information for

the phenology analysis. The tests were performed by comparing the phenological transition dates extracted from the image time series to the corresponding visual estimates, and to those observed in the field in the frame of phenological observation network of Finland, which covers a long latitudinal transect ranging from 60°N to nearly 70°N (Poikolainen et al., 1996; Pudas et al., 2008).



## 2 Materials and Methods

### 2.1 Sites and camera installations

Camera sites cover nearly the full range of climatic variations observed in Finland, their location ranging from the hemiboreal Tvärminne to the sub-arctic Kaamanen (Figure 1, Table 1). Three of the northern sites are wetlands [Sodankylä wetland, Kaamanen, Lompolojänkkä Integrated Carbon Observation System (ICOS) sites] and two are dominated by *P. sylvestris* (Scots Pine)(Sodankylä ICOS site, Värriö), and one by *Picea abies* L. Karst (Norway spruce) (Kenttäröva, ICOS site). The Paljakka site in central Finland is dominated by spruce and it belongs to the long-term phenology monitoring network of Luke, as does the mixed species site Parkano in southern Finland. The other southern sites are dominated by *P. sylvestris* (Hyytiälä ICOS site), *Picea abies* L. Karst (Punkaharju, Tammela Level II monitoring sites of the International Co-operative Programme on Assessment and Monitoring of Air Pollution Effects on Forests, ICP), or have mixed or deciduous coverage (Tvärminne, Lammi Long Term Ecosystem Research (LTER) sites). The sites vary in their ancillary measurements, the most intensively measured sites being the ICOS sites in Hyytiälä and Sodankylä while the Suonenjoki *P. sylvestris* site only hosts a meteorological station.

Table 1 Cameras and sites used in this study, the total numbers of pixels analysed for each site, and DOI to original image time series. Coordinates are in decimal degrees WGS84.

No.	Site	Lat.	Lon.	Camera view / dominant species	Species	Spring period s in data	Autum n period in data	Pixels in target ROI	DOI <sup>2</sup>
1	Hyytiälä (crown)	61.85	24.30	Forest canopy / <i>P. sylvestris</i>	<i>B. pendula</i>	2014-2016	2014-2016	26695 <sup>1</sup>	10.5281/zenodo.815559
2	Kaamanen	69.14	27.27	Wetland / <i>Sphagnum</i> spp.	<i>B. pubescens</i>	2015-2016	2015 – (2016)	39527	10.5281/zenodo.815553
3	Kenttäröva (canopy)	67.99	24.24	Forest canopy / <i>P. abies</i>	<i>B. pubescens</i>	2015-2016	2015-2016	51370	10.5281/zenodo.815519
4	Lammi	61.0	25.0	Mixed	<i>B.</i>	2016	2016	59865	10.5281/zenodo.81554

	(landscape)	5	4	landscape / <i>B. pendula</i>	<i>pendula</i>			0	2
5	Lompolojännkä	69.80	24.21	Wetland / grasses	<i>B. pubescens</i>	2015	2015	324728	10.5281/zenodo.815555
6	Paljakka	64.68	64.68	Mixed landscape / <i>P. abies</i>	<i>B. pubescens</i>	2016	2015	16867	10.5281/zenodo.815529
7	Parkano	62.03	23.04	Mixed landscape / <i>B. pendula</i>	<i>B. pendula</i>	2016	2015-2016	487184	10.5281/zenodo.815487
8	Sodankylä, wetland	67.37	26.65	Wetland / <i>Sphagnum</i> spp.	<i>B. pubescens</i>	2014-2015	2014-2016	64076	10.5281/zenodo.815485
9	Suonenjoki	62.64	27.05	Forest crown level / <i>P. sylvestris</i>	<i>B. pendula</i>	2016	2015-2016	155758	10.5281/zenodo.815489
10	Tammela (canopy)	60.65	23.81	Forest canopy / <i>P. abies</i>	<i>B. pendula</i>	2014-2016	2014-2016	46022	10.5281/zenodo.815450
11	Tvärminne	59.84	23.25	Mixed landscape / <i>P. sylvestris</i>	<i>B. pendula</i>	2016	2016	77705	10.5281/zenodo.815550
12	Värriö (crown)	67.75	29.61	Forest crown level / <i>P. abies</i>	<i>B. pubescens</i>	2015-2016	2016	191423	10.5281/zenodo.815534
	N site-years					20	21		

149 1 Hyytiälä crown camera had resolution 1024 x 768 while others had 2594 x 1944.

150 2 Peltoniemi et al., 2017

151

152 All cameras are set to a fixed white balance, quarter of the maximum resolution (5 MPix), targeted

153 northwards where feasible and triggered for half-hourly submission of snapshots to an ftp server,

154 excluding the night hours. All of the sites and analyses of this study used image time series acquired

155 with StarDot NetCam SC5 cameras.

156

## 2.2 Phenological analyses

### *Material for phenological analyses*

In this study, we used 12 networked cameras for the analyses of spring and autumn phenology of *Betula* spp from 2014-2016. Installation and operation dates of the cameras varied, and not every camera covered the whole period. Some cameras had suffered from damages or mounting failures, and had been reinstalled causing gaps in time series. Therefore, depending on the availability of images at the camera site during this period, we analysed the phenology of either a full year or a limited spring or autumn period, as indicated in Table 1.

The ROIs were selected subjectively to cover the sub-regions that best represented the birch crowns within the camera view (Appendix A). Consequently, ROIs varied in shape, size and the number of sub-polygons defining the ROI, depending on the features and number of suitable targets. The number of sub-polygons varied from 1 to 6 per image time series, the ROIs most often representing individual trees in the images.

Targeted crowns in the ROI also had a variable background, depending on whether there were conifer crowns, peatland vegetation or sky in the background of the targeted crown. Some of the targets were large and had a uniform background, while others were small and had uniform background. Based on preliminary analysis, we excluded two targets that had seedling birches against an understory vegetation background, to avoid risk confusing two distinctive sources of vegetation signals, and one target with a single distant birch tree against conifers (excluded partly due to camera movement). The remaining unclear cases were analysed in order to study if their phenological transition dates are plausible with respect to other sites' results and other materials. For these purposes, targets were classified either as distant-small and near-clear, and effects of category tested in Anova (see below).

For each ROI, we calculated the mean green chromatic coordinate (GCC) and red chromatic coordinate (RCC) as  $GCC = G / (R + G + B)$  and  $RCC = R / (R + G + B)$ , where R, G, and B, and pixel red, green, and blue channel digital numbers, respectively. For the extraction of digital numbers from images, and the calculation of these indices, we used a custom made program (FMIPROT, Tanis et al., submitted).

The pixels of ROIs with poor or excess exposure were excluded from the GCC calculation; we used only pixels with digital numbers between 30 and 254, to avoid too dim and overexposed pixels and their non-linear effects on GCC and RCC. The use of R, G and B threshold therefore eliminated images from the darkest periods in winter in northern Finland.

The image time series consisted of half-hourly images that were taken within the daily period of 8:00-16:00 UTC+2. From all images available for a day, and for all days, we calculated daily medians of GCC and RCC, which were used in subsequent analyses. We also calculated daily 90<sup>th</sup> upper percentiles, but as our preliminary analysis showed that the median provided less noisy (but otherwise very similar) results, we used the daily medians in the final analyses.

#### *Turning point estimation from continuous color indexes: phenological transition dates*

We fitted continuous curves to the GCC and RCC data (daily medians), which allowed the estimation of turning points that correspond to the transition dates. The curve fits were made between DOY 90 (Mar 1<sup>st</sup>) and 310 (Oct 6<sup>th</sup>), except for the northernmost Kaamanen site, where we started the fits on DOY 125 (Mar 25<sup>th</sup>), and the second northernmost Värriö site, where we ended the fit period on DOY 300 (Sep 27<sup>th</sup>). Focusing the fits on these periods eliminated the infection of colour signals by canopy snow cover

and dark winter days, which can bias the signal and add variation to GCC (Linkosalmi et al., 2016), and thus negatively influence the curve fitting and subsequent extraction of GCC transition dates. Any gaps in the GCC time series, due to low light or camera malfunctioning, were filled with linear interpolation.

We fitted different versions of double logistic functions (Gu et al., 2009, Elmore et al., 2012, Klosterman et al., 2014) to the GCC data. The formulation by Gu et al. (2009) produced curves that systematically fitted well to the data. While the methods of Klosterman et al. (2014) and Elmore et al. (2012) typically produced good fits, but sometimes they produced very poor fits. This was likely because Klosterman et al. (2014) and Elmore et al. (2012) included more parameters than Gu et al. (2009), which made the regression more unstable. Therefore in subsequent analyses, we only used the method of Gu et al. (2009) as implemented in *phenopix* (Filippa et al., 2016), which fits the function of the following form:

$$GCC(t) = y_0 + \frac{a_1}{\left[1 + e^{(-b_1^{-1}(t-t_{01}))}\right]^{c_1}} - \frac{a_2}{\left[1 + e^{(-b_2^{-1}(t-t_{02}))}\right]^{c_2}} \quad (1)$$

where  $GCC(t)$  is the GCC median of day  $t$ , and  $y_0$ ,  $a_1$ ,  $b_1$ ,  $c_1$ ,  $t_{01}$ ,  $a_2$ ,  $b_2$ ,  $c_2$ , and  $t_{02}$  are parameters to be estimated. The equation is composed of two modified logistic functions that characterize increasing and decreasing parts of the season. The function is flexible, but may omit finer variation of season progression. The uncertainty of fits and subsequent estimation of transition date estimates (see below) was made with a method implemented in `GuFit()` function of the *phenopix* library, i.e. by repeating the fit 100 times by introducing uncertainty to the observations.

Statistical performance of the double exponential fits was evaluated and compared by calculating the root mean squared deviation (RMSD) by site and year as  $RMSD = (\text{sum}((g_i - p_i)/n))^{1/2}$  where  $i$  is day,  $g$  is the observed GCC,  $p$  is the modelled GCC, and  $n$  is the number of days. The median and the 2.5 and 97.5

percentiles of RMSD were estimated for each site-year combination from the ensemble of 100 fits so as to evaluate the uncertainty of fits.

We also used *SplineFit()* function from the *phenopix* R library (Filippa et al., 2016) to fit cubic spline regressions to the daily time series of GCC and RCC (Filippa et al., 2016). The fit is sensitive to the selection of the degree of freedom of the fit (parameter *k*). We allowed the fit algorithm to select the optimum *k* automatically. We also made preliminary tests of the effects of *k* selection to the fits and subsequent transition date estimation, but found out that the varied *k* values around the optimal *k* produced very similar estimates of season transition dates, and hence report only the fits and transition dates with the optimal *k*. RMSD was estimated similarly as with the double-logistic fit.

We extracted phenological transition dates from the fitted curves with the *PhenoExtract()* function of the *phenopix* library. For spline fits, we used the '*derivatives*' method that extracts the dates when the GCC increase is steepest in the spring (SOS, start of season) and when the autumnal decrease in GCC is steepest (EOS, end of season), as implemented in the *greenbrown* R library (Forkel et al., 2015, Forkel and Wutzler 2015). The date of the maximum GCC during the season was denoted as POP (peak of season). Additionally, we estimated the end of season from the peak of spline smoothed RCC (EOSr, end of season RCC). The methods for estimating the transition dates are shortly described in

Table 2. For the double logistic fit using the method presented by Gu et al. (2009), we extracted the transition dates estimating the first marks of fractional increase of green color (GCC) of leaves (UD, upturn date), the stabilization date of this fraction to summer levels (SD, stabilization date), the first marks of autumn decline (DD, downturn date), and the levelling of GCC to late autumn levels (RD, recession date).

Table 2 Automatically estimated phenological transition dates from image time series.

Acronym	Variable name	Explanation	Reference
---------	---------------	-------------	-----------

DD	Downturn date	Intersection of horizontal lines through summer GCC plateau and line tangential to the peak senescing point that is estimated from the minimum of the first derivative of GCC.	(Gu et al., 2009); (Filippa et al., 2016)
EOS	End of season (from GCC)	Midpoint of the autumn senescence period, defined as the maximum of the first derivative of the senescing curve. Used for spline fitted seasonal development.	(Forkel et al., 2015); (Forkel and Wutzler 2015)
EOSr	End of season (from RCC)	Maximum of the spline fitted RCC curve in the autumn.	-
POP	Peak of season	Maximum of the spline fitted GCC curve in the season.	(Filippa et al., 2016)
SOS	Start of season	Used for spline fitted seasonal development.	(Forkel et al., 2015); (Forkel and Wutzler 2015)
RD	Recession date	Intersection of horizontal lines through autumn minimum GCC and line tangential to the peak senescing point that is estimated from the minimum of the first derivative of GCC. Occurs generally when leaves grow fast in the spring.	(Gu et al., 2009); (Filippa et al., 2016)
SD	Stabilization date	Intersection of horizontal lines through summer GCC plateau and line tangential to the peak recovery point that is estimated from the maximum of the first derivative of GCC.	(Gu et al., 2009); (Filippa et al., 2016)
UD	Upturn date	Intersection of horizontal lines through spring minimum GCC and line tangential to the peak recovery point that is estimated from the maximum of the first derivative of GCC. Occurs generally when leaves grow fast in the spring.	(Gu et al., 2009); (Filippa et al., 2016)

#### Phenological field observations

We used a subset of data for years 2015-2016 from the phenological field observation network operating in Finland (Poikolainen et al., 1996; Pudas et al., 2008). The number of locations in this subset was 17 and the sites covered the latitudinal domain of phenology cameras (Table 3, Figure 1). At each stand the observations were made individually by observing five medium-sized and healthy birches.

Table 3 Phenological field observation sites used in the study.

ID	Site	Latitude	Longitude	Species
a	Aulanko	61.02	24.46	<i>B. pubescens</i>
b	Joensuu	62.6	29.73	<i>B. pubescens</i> & <i>B. pendula</i>
c	Kannus	63.93	23.89	<i>B. pubescens</i> & <i>B. pendula</i>
d	Kevo	69.76	27.01	<i>B. pubescens</i>
e	Kolari	67.35	23.83	<i>B. pubescens</i> & <i>B. pendula</i>

f	Lapinjarvi	60.62	26.17	<i>B. pubescens &amp; B. pendula</i>
g	Muddusjarvi	69.06	27.11	<i>B. pubescens</i>
h	Muhos	64.82	26.01	<i>B. pubescens &amp; B. pendula</i>
i	Oulanka	66.35	29.32	<i>B. pubescens &amp; B. pendula</i>
j	Parkano	62.03	23.04	<i>B. pubescens &amp; B. pendula</i>
k	Preitila	60.45	22.76	<i>B. pubescens &amp; B. pendula</i>
l	Punkaharju	61.81	29.33	<i>B. pubescens &amp; B. pendula</i>
m	Ruotsinkyla	60.36	24.99	<i>B. pubescens &amp; B. pendula</i>
n	Solbole	60.04	23.04	<i>B. pubescens &amp; B. pendula</i>
o	Suonenjoki	62.64	27.06	<i>B. pendula</i>
p	Varrio	67.75	29.61	<i>B. pubescens &amp; B. pendula</i>
q	Vesijako	61.39	25.05	<i>B. pubescens &amp; B. pendula</i>

264

265 According to the field guide of the network (Kubin et al., 2007), the budburst date ('Budburst') was  
 266 recorded when half of the leaves of *Betula* spp. have emerged from the bud. However, these leaves have  
 267 not yet unfolded, i.e. the blade and midrib of leaves are not yet visible. The guide complements this  
 268 definition by stating that at budburst the trees show the first marks of green colour in spring from a  
 269 distance. We compared these dates to images extracted UD and SOS.

270

271 An estimate of the date when leaves have grown to full size and thickness ('Leaves grown') was  
 272 recorded when there was no apparent increase in size or thickness of the individual leaves of the crown.  
 273 At this time crowns of birches usually have also reached their full density. We compared these dates to  
 274 image extracted SD.

275

276 The leaf yellowing date ('Leaves yellow') was defined as the date when 50% of individual crowns have  
 277 yellow leaves due to the normal autumnal senescence process, and not due to diseases such as leaf rust  
 278 fungi. We compared these dates to image extracted EOSr.

279

280 The leaf fall date ('Leaves fallen') was defined as the date when 50% of leaves of individual tree crowns  
 281 had shed leaves.

282



*Visual transition estimates from image time series*

Using turning point analysis we also compared the transition dates extracted from the image time series to the visually determined season transition dates from the same image time series, so as to verify how well the automated analysis is able to detect transition dates and periods determined visually by going through image time series (half-hourly images). When several targets were available from an observation site, the means of all trees of focal species were recorded to visually determine the transition dates. This also corresponds to the automatically extracted GCC that were estimated either as a mean of several polygons if several trees of same species were present, or as a mean of a wider image area of homogeneous canopy.

An experienced phenological observer, who had no other association with the data analyses, developed and used a protocol for detecting birch leaf budburst, maturation and leaf autumn colouring from image time series. The observer followed, as closely as possible, the same field guidance as was used within the phenological observation network (Kubin et al., 2007), although obvious modifications were introduced due to the low resolution of images and, in some cases, distant elements. The estimation of the budburst date of *Betula* spp. relied on the colour change of the tree crowns, as it was rarely possible to distinguish individual leaves. This definition presumably yields results that are very close to those from the observation network, which also takes note of the colour change of the canopy from a distance. We compared these budburst estimates to UD and SOS.

An image-based estimate corresponding to the 'Leaves grown' estimate of the observation network was defined as the date when the birch crowns have reached their full density. After this date the crowns start to lose their distinctive light green colour. We compared these estimates to SD and POP.

Instead of a single date corresponding to the leaf yellowing date, we estimate a period during which leaves turn yellow, so as to better quantify the correspondence with the automatically estimated DD and RD. These dates were defined as the date at which 10% and 90% of leaves are yellow or brown, and it was assumed that the field-observed date when 50% of leaves were yellow occurs between these dates. A similar approach was used to estimate the leaf fall period (10% and 90% fallen). These dates presumably contained the field-observed date (when 50% of leaves had fallen), had we had a direct field observation of the imaged trees. We also compared these dates to EOS and EOSr.

### *Statistical comparisons and effect analyses*

Irradiance conditions have been earlier found to influence GCC of conifers, rendering images useless when it is too dark (Linkosalmi et al., 2016). We studied if radiation and temperature can explain day-to-day variation in GCC data, e.g. by exposing different and a variable number of pixels for the GCC or RCC calculation of the ROI depending mostly on the light available to the inner canopy, or by influencing camera image cell sensitivity, respectively. Given that day-to-day variation of midday irradiance and temperature can be large in comparison to slower pace shifts occurring in phenology, the absence or small contribution of these variables would indicate that they are unlikely to influence the analyses of derived transition dates. Alternatively, a large effect of these variables would indicate that irradiance changes bias the transition date estimates based on image time series.

In order to estimate how large an effect light conditions and temperature can have on the GCC detected with the cameras, we fitted a linear mixed effects model with a temporal autocorrelation (AR1) term. Because GCC values are not comparable across sites, we scaled them by site and year to have mean of zero and unit standard deviation. The AR1 structure removes the trend-like variation in GCC by assuming that residuals of the model are auto-correlated, meaning that previous day's GCC is accounted for in the

prediction of the present GCC, which evidently clears the data from seasonal changes of GCC. The model formulation makes it possible to estimate the direct effects of light conditions and temperature on the present GCC. The fitted model form was

$$y_{ijt} = a_0 + a_1 T_{ijt} + a_2 G_{ijt} + a_3 G_{ijt} \times T_{ijt} + b_{0i} u_{s_i} + b_{1ij} u_{Y_{ij}} + \varphi_{ij} y_{ij(t-1)} + e_{ijt},$$

(2)

where  $y$  is the GCC observation, and subscripts  $i$ ,  $j$ , and  $t$  denote the site, year, and day of the observation, respectively. Fixed terms included the intercept ( $a_0$ ) and daily mean temperature ( $T$ ) and global radiation ( $G$ ), and interaction of  $T$  and  $G$ , with respective coefficients  $a_1$ ,  $a_2$ , and  $a_3$ . For  $T$  and  $G$  we used spatially downscaled estimates made for the nearest grid point (Venäläinen et al., 2005). The model has intercepts  $b_{0i}$  and  $b_{1ij}$  for random terms  $u$  for sites ( $s$ ) and years ( $Y$ ), respectively. Autocorrelation was modelled with AR1 process having an autocorrelation coefficient  $\varphi$ . The model was fitted with the *lme* function of the R package *nlme* (Pinheiro et al., 2015). The model was fitted to a selected subset of the data including Hyytiälä (2014-2016), Kenttäröva, Sodankylä and Tammela (2015-2016), Kaamanen (2015), and Tvärminne and Värriö and Lammi (2016), which all had full time series from UD-10 to RD+10 days.

For testing the relationship of GCC and RCC based estimates and the estimates based on visual interpretations of corresponding transition dates, we plotted 1:1 graphs, fitted linear models between the estimates and estimated their mutual correlation coefficient. We also used linear mixed effects models to estimate the significance of differences between the transition dates (UD, SOS, EOSr) estimated with different methods. Method types included GCC (or RCC) based estimation, visual

interpretation, and field observation. Field observation was not conducted at camera sites, so we also included latitude (and year) in this regression. The transition date  $y_{it}$  estimated with these methods was modelled as

$$y_{it} = a_0 + a_1 l_i + a_2 Y_{it} + a_3 l_i \times Y_{it} + a_4 E_{it} + b_{0i} u_{s_i} + e_{it} \quad (3)$$

where  $a_0$  is the intercept,  $a_1$ ,  $a_2$  and  $a_3$  are the coefficients for  $l_i$  (the latitude (easting (m) of the Finland Uniform Coordinate System) of the site and  $Y_{ij}$  (year), and their interaction, respectively.  $E_{ij}$  is the method of the observation of the observation, having  $a_4$  as its coefficient. Coefficients  $b_{0i}$  are for the random terms  $u$  for sites  $s_i$ , and  $e_{it}$  is the normally distributed error. We also estimated separate linear regressions to investigate the latitudinal relationships of transition dates obtained from cameras and the field observation network. All statistical analyses were made in R (R Core Team, 2015).

### 3 Results

#### *Fits to GCC and RCC data*

Double logistic fits replicated the upturn of the GCC response at season start for the birch ROI at Lammi (Appendix A, Figure 2) and the other sites as well as the spline regression (Appendixes A and B). However, the slope of the senescing trail of GCC after the season peak varied by site and year, and double logistic function did not always fit to shapes of the trails (Appendix B). The parametrically more flexible spline regressions were able to catch the whole seasonal course of GCC and had better RMSD than double logistic fits (0.00172 vs, 0.00216, see Appendix C). However, being sensitive to within season variation of GCC, it sometime falsely interpreted season end from GCC data (EOS) (Figure 2). For the season end date, estimates obtained for red peak (EOSr) of the spline smoothed RCC were always plausible (Figure 3, Appendix B).

For double logistic fit, the uncertainty of transition date extraction varied. Uncertainty ranges of UD, SD, and RD (10th - 90th percentile range) were on average 1.0, 1.4, and 2.5 d, respectively, while for DD it was 5.6 d.

Based on the autocorrelation model (Eq. 1), radiation had small, yet significant, effect on the observed day-to-day GCC variation (Table 4). Altogether, the fixed covariates (temperature, radiation and their interaction) were able to explain 1.6% of day-to-day variation of GCC (average across site-years).

Table 4 Results of the model (fixed terms) explaining day-to-day variation in GCC (scaled to zero mean and unit standard deviation) with daily mean temperature ( $T$ , °C) and radiation sum ( $G$ , MJ m<sup>-2</sup> day<sup>-1</sup>), and their interaction.  $\varphi$  is the autocorrelation coefficient of the AR1 autocorrelation structure.

<i>Dependent variable:</i>	
	GCC
$a_0$ (intercept)	-0.27 (0.17)

$a_1(T)$	0.0014 (0.0036)
$a_2(G)$	-0.0120*** (0.0030)
$a_3(G \times T)$	0.00068*** (0.00021)
$\varphi$	0.97
Observations	2317
Log Likelihood	-221.38
Akaike Inf. Crit.	458.77
Bayesian Inf. Crit.	504.75

---

Note: \*p<0.1; \*\*p<0.05; \*\*\*p<0.01

None of the transition dates, nor the length of green-up period (SD-UD) or season (EOSr-SOS) were explained by the type of the ROI (distant and small vs. near and clear) as analysed by linear mixed effects models with and without the latitude covariate (all  $p > 0.05$ , models not shown, for data see Appendix C).

#### *Comparison of season start estimates*

The estimates of transition dates for start of the season (SOS) were significantly related to the visually estimated budburst date (Figure 3), but they were on average 6.37 days later than the visual estimate and field observation of the budburst date (Table 5).

The upturn date (UD) was also significantly associated with the budburst date (Figure 4**Error! Reference source not found.**). The camera-observed UD was on average 3.46 and 4.08 days ahead of the visual and field observed budburst (Table 5). It is notable that the estimation of UD was occasionally interfered with by the preceding snowmelt that occurred in the background of the birch targets (Kenttäröva and Värriö), which reduced the correlation.

The visual estimates of budburst were mostly between the estimated UD and SOS dates (16 out of 19 cases where they could be compared) (Appendix C Table C.3).

#### *Comparison of season end estimates*

The autumn peak of RCC (EOSr) was clearly associated with the visual season end estimates (90% of canopy yellow) (Figure 3). EOSr also associated with the date when 90% of leaves were interpreted as fallen from the tree crowns. EOS was less clearly associated with these events than EOSr (Figure 4). We found no statistical evidence that EOSr would be different from the visually interpreted leaf yellowing date (Table 5).

DD and RD dates were weakly related to dates when 10% and 90% leaves were yellow or fallen.

#### *Comparison of midseason estimates*

POP was related with canopy maturation (leaves grown) but the relationship deteriorated with increasing DOY for the northern sites (Figure 3).

SD was correlated with the visual estimate of the date of full-sized leaves but on average was dated earlier (Figure 4).

Table 5 Model of transition dates with latitude, with method and year of observation as covariates. Yellowing date in field was recorded when 50% canopy was yellow, while visual camera based observations were made when 90% of the canopy was yellow. Interaction between the latitude and observation type were insignificant and not included in the models ( $p>0.05$ ). \*  $p<0.1$ ; \*\*  $p<0.05$ ; \*\*\*  $p<0.01$ . Note, latitudinal slope is composed of two species, *B. pubescent* in north and *B. pendula* in south.

Camera-based estimate	UD	SOS	EOSr
Field/Vis. Observation	Budburst	Budburst	Leaves yellow
Intercept (camera, 2015)	-18.43 (12.62)	-7.81 (12.82)	439.71 *** (23.52)
Latitude (km)	2.14 *** (0.18)	2.14 *** (0.18)	-2.47 *** (0.33)

Year 2016	-5.17 <sup>***</sup> (0.76)	-6.24 <sup>***</sup> (0.81)	-8.03 <sup>***</sup> (1.26)
Field obs.	4.08 <sup>***</sup> (1.26)	-6.41 <sup>***</sup> (1.29)	-1.26 (2.32)
Visual obs.	3.46 <sup>***</sup> (1.12)	-6.37 <sup>***</sup> (1.18)	1.12 (1.74)
Observations	89	90	82
Log Likelihood	- 243.77	-251.28	-263.08
Akaike Inf. Crit.	501.53	516.56	540.15
Bayesian Inf. Crit.	518.55	533.66	556.56
Note:	* p<0.1; ** p<0.05; *** p<0.01		

#### *Latitudinal gradients of transition dates*

Our camera network made it possible to determine the latitudinal gradient in transition dates across Finland. There was a clear south-north trend in the season lengths (Figure 6), the seasons becoming shorter by 4.81 ( $\pm 0.63$  Std. Err.) days per 100 km when moving northwards (Table 6). This trend was caused by a later start (both SOS and UD) and earlier end (EOSr) of the season in the north, which increased by 2.07 ( $\pm 0.42$ ; for UD 1.94  $\pm 0.40$ ) and decreased by 2.77 ( $\pm 0.52$ ; for SD 2.18 $\pm 0.49$ ) days per 100 km northwards, respectively, in year 2016. The latitudinal slopes were not statistically different between the years (Figure 7, Table 6, tested during Table 6 analyses). There were no differences between latitudinal trends of field observations and the corresponding GCC and RCC-based estimates (UD and budburst, SOS and budburst, and EOSr and yellowing; interactions were not significant, not shown). POP and RD also had significant latitudinal relationships, but EOS and SD did not (Appendix C.3). Interestingly, the residual standard errors (RSEs) of the camera derived latitudinal relationships for SOS



and UD were almost as small as those of the field observed budburst date, and EOSr and SD had even a smaller RSE than the field observed leaf fall date.

Table 6 Latitude relationships of the phenological transition dates of season. Budburst and yellowing dates are estimated from field observations in the phenological network in Finland. Others are extracted from image time series. Values in parentheses are standard errors of estimates. \* p<0.1; \*\* p<0.05; \*\*\* p<0.01. Units of dependent variables are numbers of days for EOSr-SOS and DOY for others. Note, latitudinal slopes are composed of two species, *B. pubescent* in north and *B. pendula* in south.

	Dependent variable (DOY)						
	EOSr-SOS	SOS	EOSr	UD	SD	Budburst	Yellowing
Intercept (2016)	462.73*** (44.82)	-10.41 (30.03)	453.74*** (37.08)	-8.66 (28.28)	-13.53 (35.25)	11.48 (19.05)	468.20*** (35.76)
Lat. (100 km)	-4.81*** (0.63)	2.07*** (0.42)	-2.77*** (0.52)	1.94*** (0.40)	2.18*** (0.49)	1.69*** (0.27)	-3.01*** (0.50)
2015	37.38 (71.44)	20.08 (47.07)	27.44 (57.53)	-22.38 (47.23)	-24.77 (53.63)	-36.01 (26.13)	-52.38 (48.57)
Lat:2015	-0.56 (0.99)	-0.13 (0.65)	-0.27 (0.80)	0.38 (0.65)	0.53 (0.75)	0.59 (0.37)	0.86 (0.69)
Observations	17	19	19	18	20	52	43
R <sup>2</sup>	0.90	0.83	0.79	0.82	0.83	0.74	0.65
Adjusted R <sup>2</sup>	0.87	0.80	0.75	0.78	0.80	0.72	0.63
Residual Std. Error	7.51	5.04	6.21	4.74	5.91	4.51	7.72
F Statistic	38.23***	24.99***	18.76***	20.67***	26.66***	45.07***	24.59***

Note: \* p<0.1; \*\* p<0.05; \*\*\* p<0.01

## 4 Discussion

We used our newly established digital camera network (Peltoniemi et al., 2017) for assessing birch phenology along a latitudinal transect across Finland. We showed that the network is useful for monitoring birch phenology, although the site-specific analyses often relied on appearances of small birches among other vegetation and in different types of positions and environments. This stems from the fact that birch phenological colour changes were distinctive enough to be discerned from background and sometimes small targets.

The networked cameras were particularly useful for detecting the start of the growing season green-up and the autumnal leaf yellowing. According to our results, UD is able to capture even minor changes in crown greenness and to date those reliably, on average, UD estimates season start less than 4 days earlier than the budburst date recorded in the field. This is close to what has been earlier observed for ash and beech in Switzerland (Ahrends et al., 2008) and for 13 deciduous sites in eastern North America (Klosterman et al., 2014). On the other hand, SOS systematically dates the season start approximately 6 days later than the visual and field-based observations and thus provides a reasonable late estimate, suggesting that the season start should be calculated as the mean of the UD and SOS dates. For typical seasonal paths of GCC, the extraction of these dates seems to be fairly insensitive to day-to-day variation of GCC.

Season end was best estimated with the RCC peak (EOSr) in the autumn, which compared well with the visually interpreted dates of season end. In practice, we showed no bias of EOSr relative to the date when 90% leaves were yellow. The indices based on GCC (DD, RD, EOS) turned out to be useless for the prediction of season end.

486

487 There are various sources of uncertainty in the analyses. Generally, the uncertainties of curves depicting  
488 the season progression, and extractions of key transition dates were surprisingly small. For double  
489 logistic fits, considerable uncertainties were quantified for a few cases where the automated extraction  
490 of transition dates were challenging due to the nature of the analysed GCC time series [Appendix B, due  
491 to unexplained variation of GCC (for SD in Sodankylä 2014), slowly declining season (for DD and or RD:  
492 Hyytiälä, Kenttäröva 2016, Paljakka 2015, Suonenjoki 2015 and 2016), or limited availability of data in  
493 spring (for SD: Paljakka 2016, Suonenjoki 2015)]. Otherwise, the typical daily mean GCC data seems to  
494 well support the extraction of transition dates, and considerable uncertainties seem to exist elsewhere  
495 than in curve fits and date extractions. This applies also to the spline fits whereby date extractions at  
496 least partially require subjective selection of smoothing parameter.

497

498 Challenges in discerning spring dates may also arise when conditions in the background of canopies  
499 change considerably during or before budburst. Snow melt in the background of target trees appears to  
500 be the most important factor in this region for biasing the season start estimates. For Värriö site UD  
501 occurred earlier (11.5-13.3 d, depending on year) than the visual observation, which is more than the  
502 average lag across sites (3.5 d). However, for SOS there seemed to be no significant snow induced bias  
503 (Appendix C). For further improvement of extraction algorithms, snow cover changes in the background  
504 of targets should be accounted for, either by more careful selection of the targets or by introducing a  
505 snow detection algorithm (Salvatori et al., 2011, Garvelmann et al., 2013; Arslan et al., 2017). Use of  
506 cameras in conjunction with high-resolution satellite imagery for the detection of the presence of snow  
507 in the vicinity of a camera could resolve some of the challenges. For instance, Sentinel-2 MSI provides an  
508 excellent data source for this and we have an already implemented method (SCAmod, Metsämäki et al.,  
509 2015) for deriving information about snow cover in 10m spatial resolution. Use of far and near remote

sensing methods together would provide spatially) and temporally complementary material, and could yield better performing snow and seasonal indices..

In our case, conifer background of birch targets had a minor effect on the GCC signal, as GCC and the extracted transition dates were in good accord with the transition dates that were visually assessed from the image time series (without any prior knowledge of GCC development) (Suppl. B). During the budburst of birches, the vegetation in the image background is rarely active, as conifers are not yet developing new shoots, which makes it easy to distinguish the GCC changes due to birch leaf growth. Co-occurring GCC changes of conifers (Wingate et al., 2015) potentially associated with photosynthetic and/or pigment recovery of conifer seem to be small in comparison to GCC changes caused by birch leaf growth. Possibly, phenological date extraction of midseason phenological phases could benefit from incorporation of background subtraction using mixing models (Keenan et al., 2013).

For colour index analyses, the small image elements were sufficient. However, small movement of the cameras at Sodankylä wetland site caused particularly uncertain transition date estimates during 2014 (Suppl. B), although the resulting transition date estimates did not differ from the main trends in other data. The images omitted from the analysis contained a site where unexpected movement of the camera mast would have obscured the analysis of small and distant targets. These problems may be partially due to defining too narrow ROI or ROI with too small safety margin to other targets, which exposes the view to variable set of pixels as the targets move due to wind, and in the longer term due to growth. Distance from camera to target, on the other hand, tends to even out colour differences. In our work, we did not find clear effects of colours of distant elements being obscured to the extent it would have hampered the analysis, but it could be problematic in locations with more frequent occurrence of excess condensation of air humidity moisture or fogs.

Other simultaneous phenological events in trees could potentially confound the season start estimation from images. Flowering of birch occurs nearly simultaneously with budburst, with some variation in the timing between individual birch trees. As a typical masting tree species, the annual variation of the amount of male and female flowers of birches can be enormous. In a good flowering year, usually those with high May temperatures, the amount of catkins is usually several thousands in a single birch (Ranta et al. 2008, Zamorano et al. 2016). Birch flowering appears brownish and blurry in distant images. GCC changes due to increased brown tones likely remain small, but large numbers of flowers during masting years may render small changes of GCC more difficult to discern, and thus could cause a small delay of UD estimates on those years. For some other species, however, flowering may cause more problems, and the use of SOS could be a better option.

Comparisons between cameras and field data are also complicated by uncertainties of field observations. We did not have camera-observed estimates of phenology from the same trees and years as the ones available from the phenological observation network, and therefore we compared different datasets when investigating the latitudinal gradient. Therefore, it is important to understand how birches in individual sites represent the other birches in the same latitude. Clearly, the best camera-based transition date estimates (UD, EOSr) were within the variation in the dates estimated from the field observations (Figure 7), although on average a few days earlier and later, respectively. Siljamo et al. (2008) found out that individual sites represent *Betula* spp. season start dates with an accuracy of 3-8 days as compared to the regional means, with northern locations such as Finland having smaller uncertainties. At a single site, our field observations showed that the budburst date varied among tree individuals of the same *Betula* spp. by 1.1 days (average of site std. dev. among individuals) and the yellowing and fall dates by 3 days within the same year. Obviously, such differences are partly driven by genetic variability among individuals (Rousi and Pusenius, 2005) and partly due to variation in the growth environment of the trees, for example trees in open environments or surrounded by other

deciduous trees experience earlier budburst in comparison to trees among conifers due to higher sun exposure. Some of the field observations were made very close to the cameras (at Parkano, Paljakka, Värriö, Hyytiälä and Suonenjoki), suggesting that the uncertainties at these sites are smaller than that estimated by Siljamo et al. (2008), likely closer to the lower 3 day boundary of Siljamo et al., 2008. Uncertainties due to field observations themselves are hard to assess, as the visual assessment is subjective, albeit informed by guidelines and conducted by experienced personnel. It should also be noted that the visual interpretation by expert observers from image time series gave similar results to field observations, which implies that the uncertainties of subjective observations may not be critical for the conclusions of our study about the usefulness of UD, SOS and EOSr for phenological monitoring. Further investigations on how well transition dates extracted from image time series represent the definitions used in particular observation networks obviously require data collection from the same trees as monitored for phenology.

Analyses of season stabilization (SD) were less successful than those of UD, SOS, and EOSr, and the deviations from the corresponding visual estimate (leaves grown) were larger, as also others have found (Klosterman et al., 2014). This may be partly due to the less precise observation of the date when leaves were grown to their full size in the field and more difficult interpretation of the image time series. GCC also saturate immediately when leaves cover image region, and further changes are not discernible although leaves still grow. Earlier studies investigating the relationship between leaf traits and GCC have found a decoupling of GCC from other leaf traits (Keenan et al., 2014; Yang et al. 2014). Keenan et al. (2014) explained GCC development, showing that GCC becomes insensitive to LAI increases at high LAI, and thus LAI peaks later than GCC. Leaves also change color during their development. During some years at some sites we observed a pronounced peak of GCC after the leaf unfolding and maturation period, which quickly declined to summer level and further but gradually decreased towards autumn, implying that methods need elaboration to account for the unexpected behaviour. This behaviour

obviously cannot be properly captured by the double exponential fit, which assumed steady decline of GCC until DD. Elaboration of methods further could benefit from a priori information about plausible season end range. The causes of this peak are unclear, but it could be related to the distinctive light green colour of new leaves, which is subsequently lost with increasing chlorophyll packing to the leaves along with albedo decrease within high chlorophyll contents (Bray et al., 1966), and possibly due to changes of leaf surface due to aging. Yang et al. (2014) observed that the chlorophyll peak lags 20 days behind the GCC peak in white oaks, which could explain the quick decline after the peak. The reasons for the variable GCC response among the sites and years remain unclear. In previous studies, variants of the double exponential fits (Elmore et al., 2014; Klosterman et al., 2014) occasionally caught the summer peak appropriately but then failed for other cases, which could be partially related to this issue.

Reflectance analyses, such as those conducted with cameras, are sensitive to the spectral distribution of the exposing radiation. A humped GCC pattern peaking at midday detected by Ahrends et al (2008), may be an indication of temporal differences in exposing light angle and colour, but it may be due to a low light exposure of objects that may render GCC analyses unreliable under low light (Sonnentag et al., 2012). Linkosalmi et al. (2016) excluded the dark winter days from the image data series analysed when they analysed an image time series of Scots pine (10.5281/zenodo.815481) and wetland (10.5281/zenodo.815485) north from Arctic Circle. Our results, on the other hand, were preconditioned by the exclusion of pixels with a poor exposure of any of the colour channels (digital number < 30), and by limiting the images used for the period between 08:00-16:00. Due to this filter, some of the darkest days were completely excluded at the northernmost sites. Our filter additionally causes the pixel number within ROI to vary with illumination, but this did not seem to be a significant factor. Therefore, we consider that our filtering cleaned the GCC time series from (any potential) systematic variation in GCC, leaving only residual variation that is too small to bias the image analysis and can be partly explained by day-to-day variability in light conditions, e.g. due to cloud cover.

## *Conclusions*

Our results extend the earlier conclusions that the camera-based phenology analysis provides a sound method for quantitative monitoring of phenology (e.g. Richardson, et al., 2007; Ahrends et al., 2008). We used networked cameras from a long latitudinal transect to study the phenology of the most widely spread deciduous species in the boreal zone, and showed that the cameras and targets provide reliable predictions of the seasonal development of birches in various conditions, particularly for season start using GCC and season end when using RCC. Small image elements were useful for the analyses, but analyses should account for their movement in wind, and in the longer term analyses, also tracking of their growth.

Moreover, the established network, together with the image analysis methods adopted, provides a good basis for automated monitoring of key phenological events for birch, which could reduce the costs of field monitoring. Season start and end dates could also be informative for the forcing of carbon balance models and in the calibration of phenology models, while we expect midseason transition dates to be much harder to use. Further research is required to understand how the species-specific phenological transitions are reflected in remote sensing phenology products (which typically aggregate signals over wide areas) in spring including nearly simultaneously occurring snow cover changes in northernmost areas. We consider that networks of modest density such as our network can cover the phenology of a few dominant tree species in the region, and can thus provide a good basis for the monitoring of species-specific phenology in the area.



## 5 Acknowledgements

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## 6 Appendixes

A. Regions of interest of image time series used in the phenological analyses of birch species.

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B. GCC time series, curve fits, and estimates of phenological transition dates (camera-based and visually interpreted).

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C. Data summary table and additional models of phenological transition dates.

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## 8 Figure captions

Figure 1 Locations of sites and cameras and phenological field monitoring plots used in the study.

Figure 2 Phenological data from Lammi site (for other sites, see Suppl. B). Top panel: horizontal black lines refer to visually determined phenological periods from image time periods. Bottom panel: radiation and air Temperature (T) are 5 day running means. Temperature sums are calculated from daily mean T with 0 °C threshold (starting from 21<sup>st</sup> Mar), and with 5 °C threshold (from 1<sup>st</sup> Jan).

Figure 3 Comparison of phenological transition dates. X-axes values (see Table 2) were extracted automatically from image time series based on cubic spline fits and y-axes values from the same plots were based on the visual examination of image time series. The solid line is the fitted regression line, and dash-dot line is 1:1 line. Each site has 1-3 observation years (indicated by different colors). Panel d: gray symbols indicate visual observation of 10% leaves fallen.

Figure 4 Comparison of phenological transition dates estimated based on double exponential fits (Gu et al., 2009) to GCC (X-axis). See Figure 3 for legends and other explanations.

Figure 5 Comparison of phenological transition dates for end of season dates based on GCC (EOS) with the visual estimates. See Figure 3 for legends and other explanations.

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850

851 Figure 6 Fitted splines and extracted phenological transition dates in a South to North (top to bottom)  
852 gradient in Finland. SOS dates (upwards triangles,  $\Delta$ ) were determined from GCC spline fits and EOSr  
853 dates (downwards triangle,  $\nabla$ ) from peaks of autumn RCC.

854

855

856 Figure 7 Latitudinal gradients in key phenological dates and measurements of corresponding variables in  
857 the field. Regression functions are in Table 6. Year did not significantly interact with latitude in the  
858 models, indicating no slope differences between the years.

Figure 1  
[Click here to download high resolution image](#)

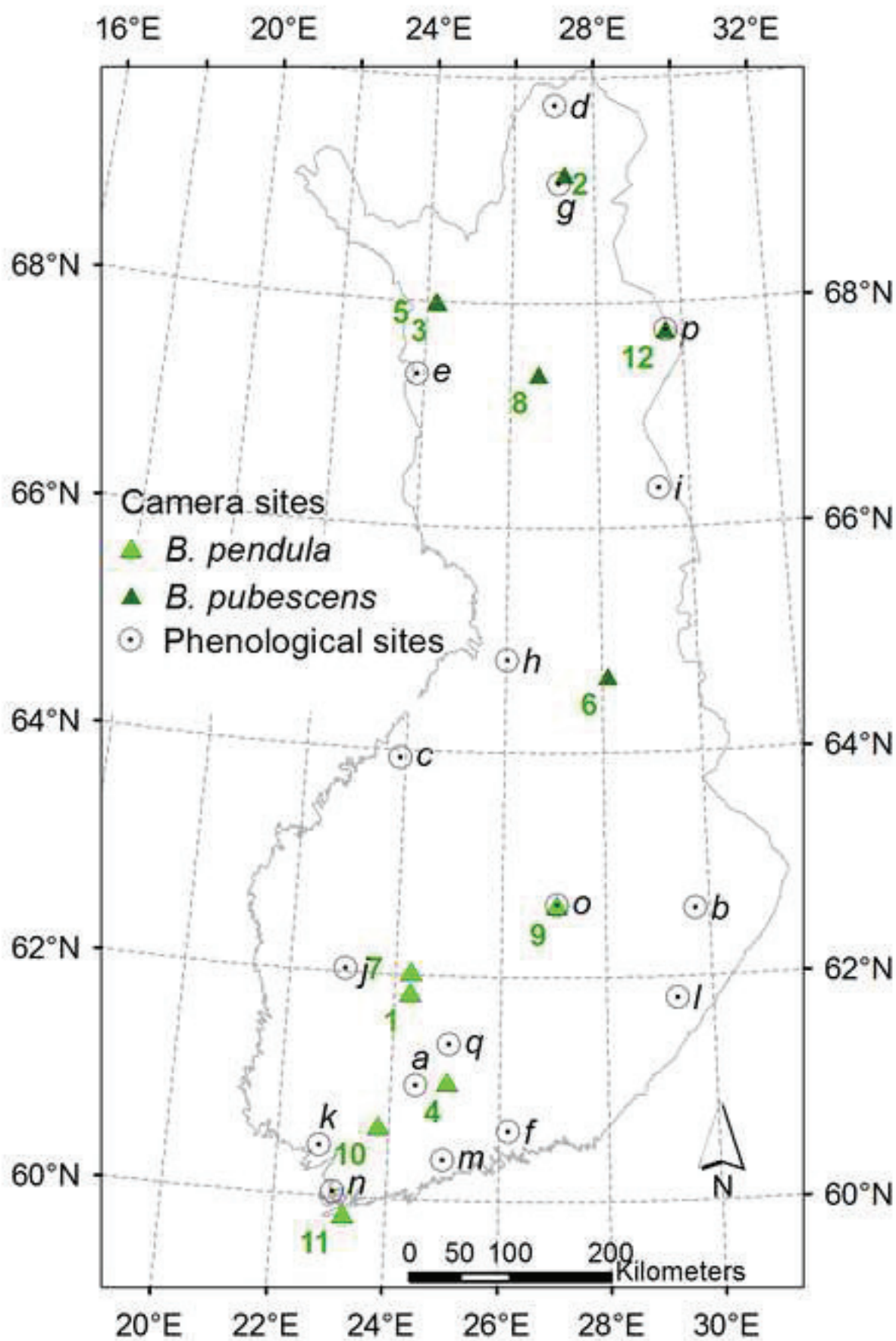
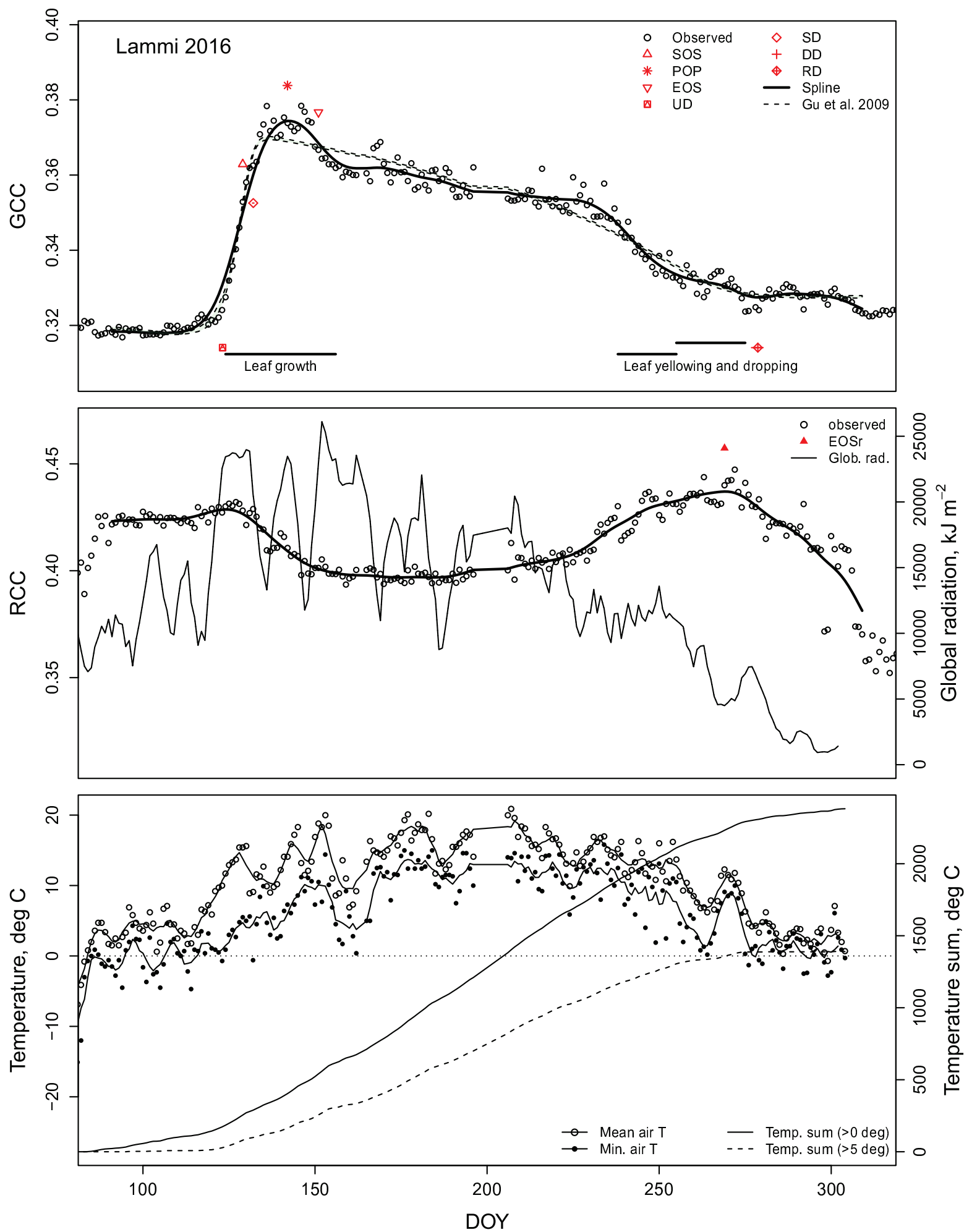


Figure 2



**Figure 3**

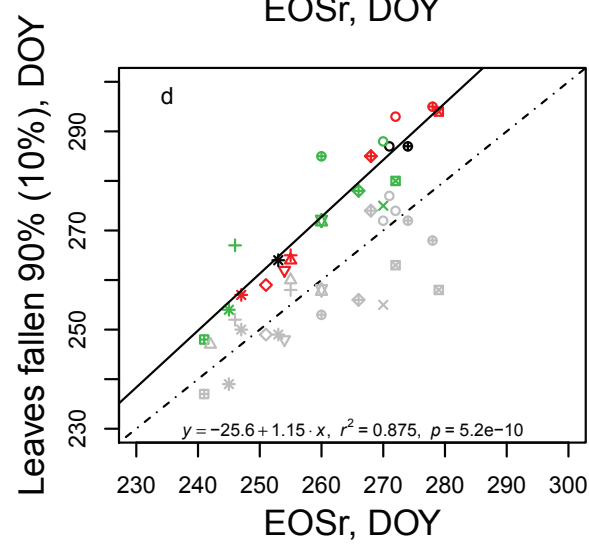
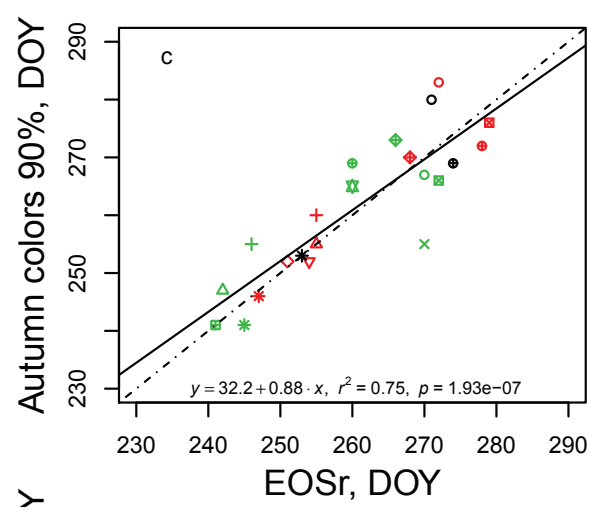
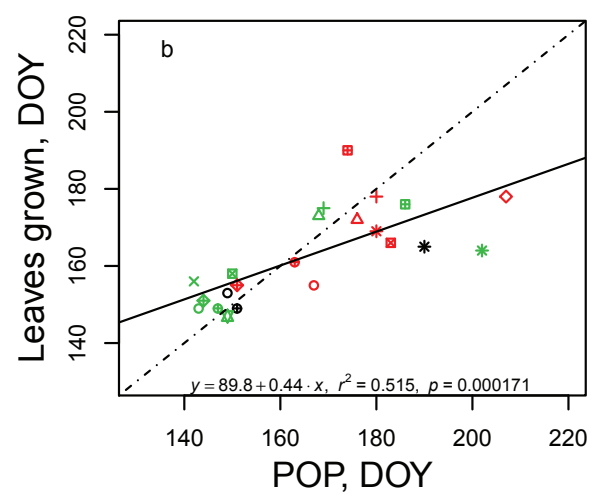
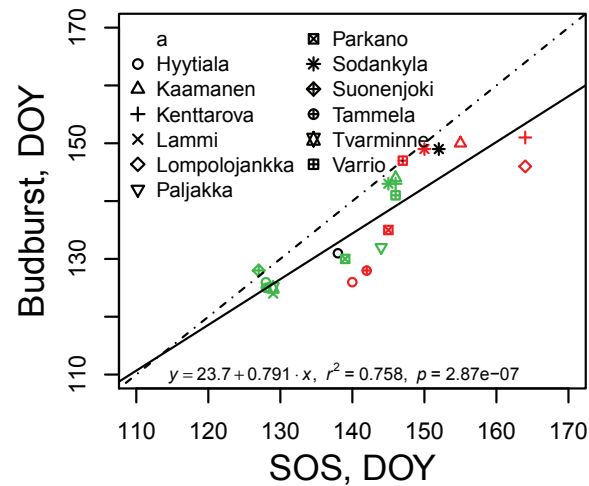


Figure 4

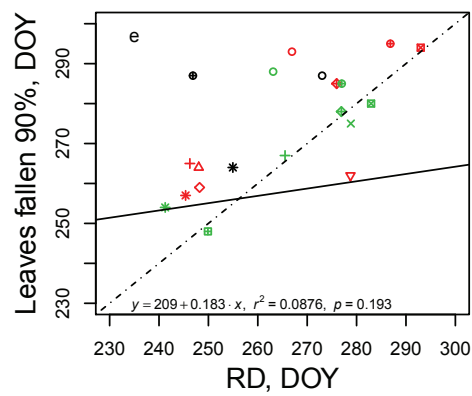
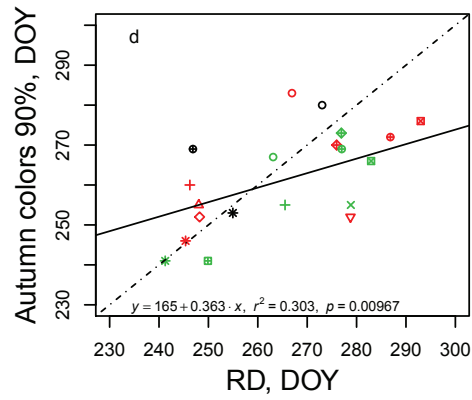
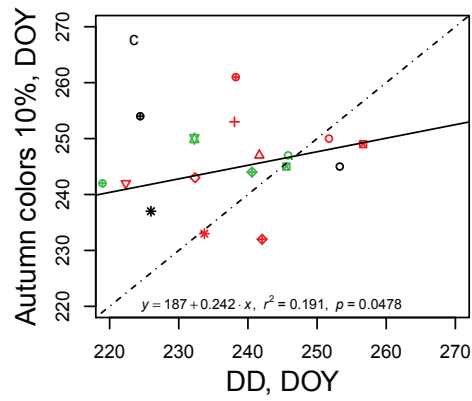
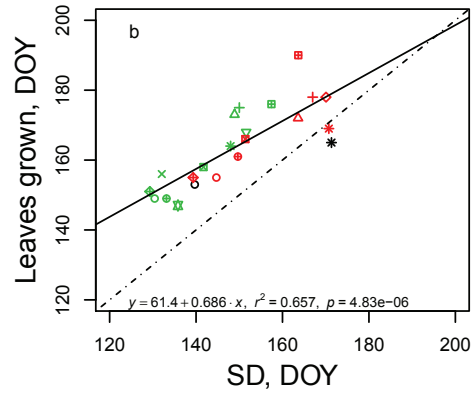
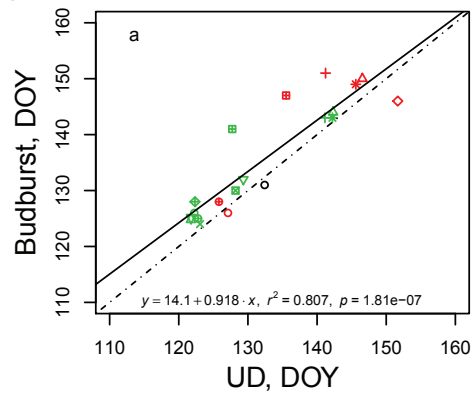




Figure 5

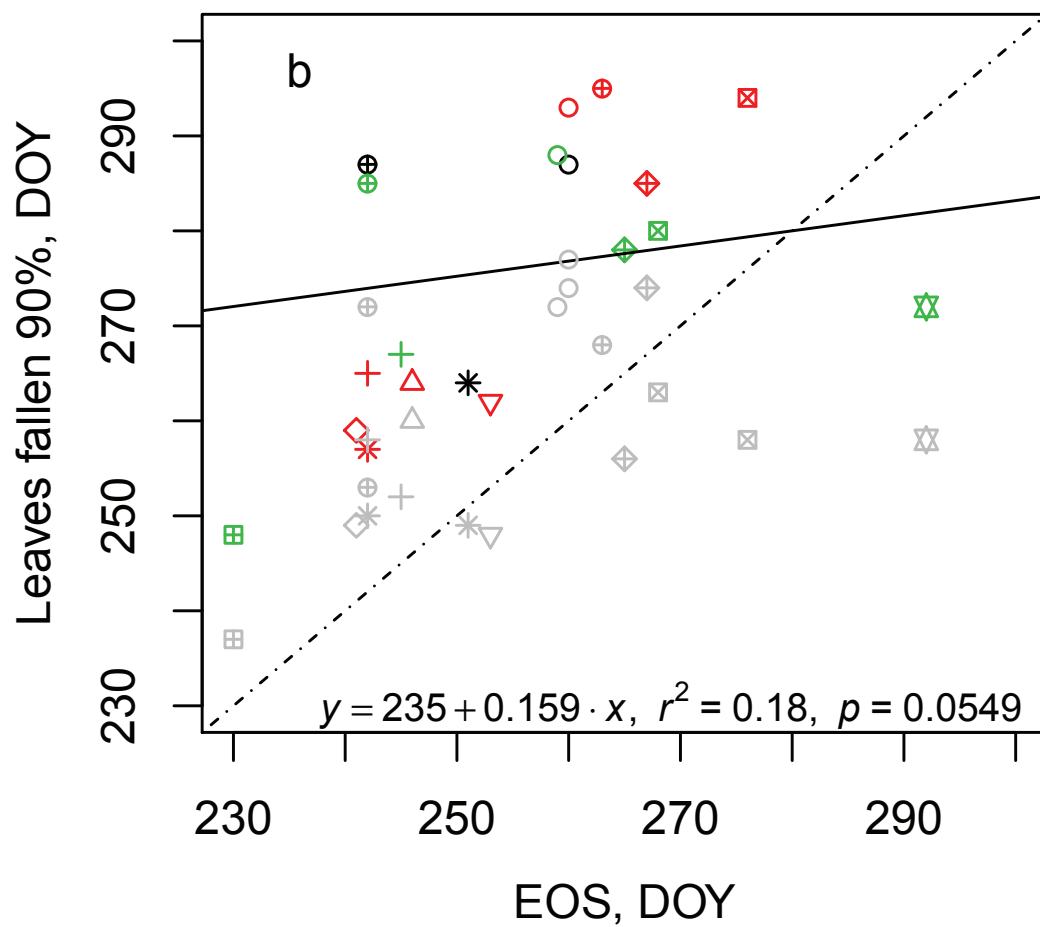
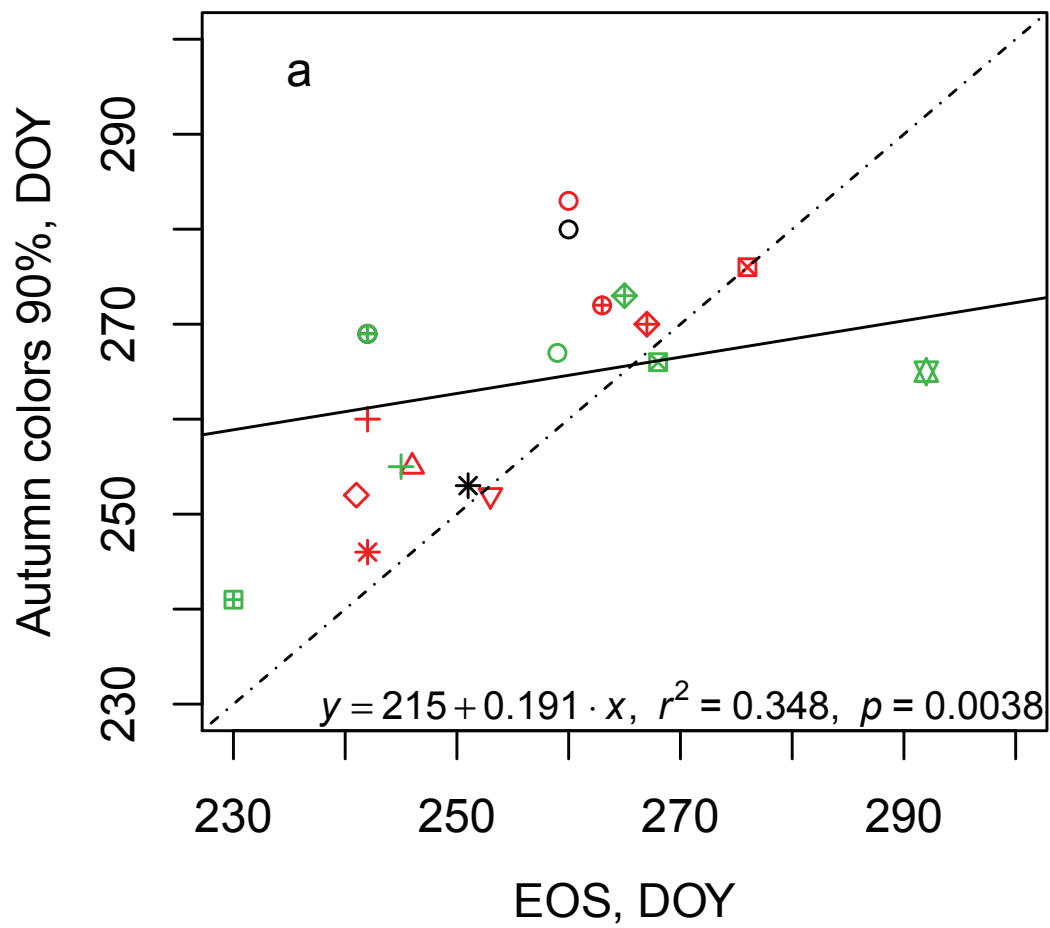


Figure 6

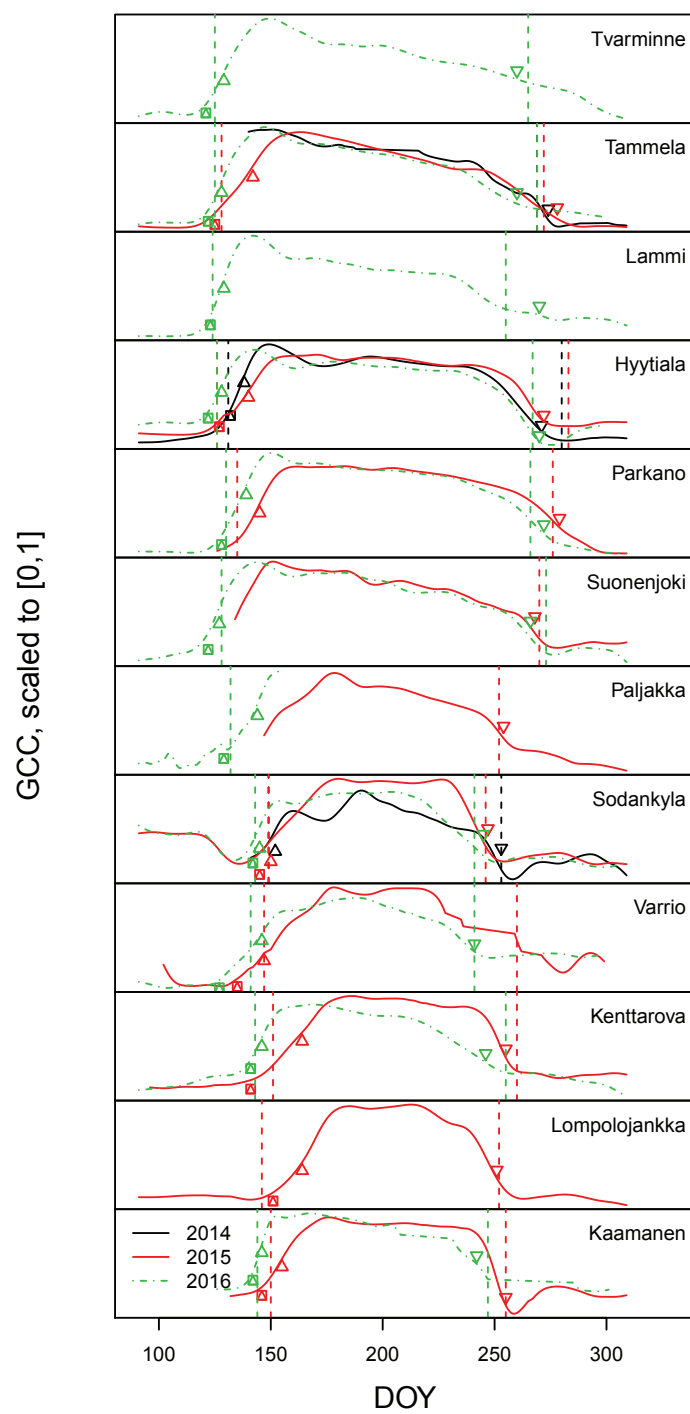


Figure 7

